IN THE CLAIMS:

Please amend Claims 1, 2, and 9 as shown below.

- 1. (Currently Amended) A method for identifying a base sequence present in a target single-stranded nucleic acid comprising the steps of:
- (a) preparing a probe array in which single-stranded nucleic acid probes of No. 1 to No. n ($n \ge 2$) are arranged as isolated spots on a substrate;
- (b) reacting a single-stranded nucleic acid, which has a base sequence fully complementary to a base sequence of one of the single-stranded nucleic acid probes and is fluorescence-labeled, with the probe array under such conditions that single-stranded nucleic acids complementary to each other form a double-stranded nucleic acid;

removing the unreacted labeled single-stranded nucleic acid; and
measuring fluorescence intensity of each spot of the probe array to obtain a
first image template pattern showing a relationship between location of the probes and
fluorescent characteristics;

- (c) performing the same operation as the step (b) for each of remaining single-stranded nucleic acid probes using a second to a nth single-stranded nucleic acid, and obtaining image template patterns of No. 2 to No. n showing a relationship between location and fluorescent characteristics of the probes;
- (d) performing the same operation as the step (b) using a sample containing the target single-stranded nucleic acid of the base sequence to obtain a sample pattern showing a relationship between a position and fluorescent characteristics;

- (e) comparing the sample pattern obtained in the step (d) with a plurality of image template patterns, the plurality of image template patterns comprising the first image template pattern and the image template patterns of No. 2 to No. n, to find an image n pieces of said image template patterns obtained in the steps (b) and (c), to find a template pattern substantially identical to the sample pattern; and
- (f) identifying the base sequence of the single-stranded nucleic acid used for the preparation of the identified image template pattern found in the step (e) as the base sequence of the target single-stranded nucleic acid.
- 2. (Currently Amended) A method for identifying a base sequence present in a target single-stranded nucleic acid comprising the steps of:
- (a) preparing a probe array in which single-stranded nucleic acid probes of No. 1 to No. n ($n \ge 2$) are No. 1 to No. n ($n \ge 3$) are arranged as isolated spots on a substrate;
- (b) reacting a single-stranded nucleic acid which has a base sequence fully complementary to a base sequence of one of the single-stranded nucleic acid probes and is fluorescence-labeled, with the probe array under such conditions that single-stranded nucleic acids complementary to each other form a double-stranded nucleic acid;

removing the unreacted labeled single-stranded nucleic acid; and
measuring fluorescence intensity of each spot of the probe array to obtain a
first template pattern showing a relationship between location of the probes and fluorescent
characteristics;

- (c) analyzing the first template pattern to locate probes and to calculate a mean value of fluorescence intensities (Fi) of the double-stranded nucleic acids having i of mismatched base pairs, where i is an integer not less than 1;
- (d) calculating a difference (F1, 0) between the fluorescence intensity of the fully complementary double-stranded nucleic acid without mismatch (F0) and the mean value of the fluorescence intensities of the double-stranded nucleic acids having one-base mismatch (F1), further calculating a difference (Fi+1, i) between a fluorescence intensity of a double-stranded nucleic acid having (i+1) base mismatches (Fi+1) and a fluorescence intensity of a double-stranded nucleic acid having i-base mismatches (Fi), and identifying i being Fi+1, i << Fi, i-1;
- (e) preparing a second template pattern of positive probe spots of probes having base sequences differing from the base sequence of the second probe by i or less bases where i is determined in said step (d), wherein negative probe spots are probes having base sequences differing from the second probe by more than i bases;
- (f) performing the same operation as the step (e) for each of remaining single-stranded nucleic acid probes and obtaining template patterns of No. 3 to No. n showing a relationship between location and fluorescent characteristics of the probes;
- (g) performing the same operation as the step (b) using a sample containing the target single-stranded nucleic acid of the base sequence to obtain a sample pattern showing a relationship between a position and fluorescent characteristics;
- (h) comparing the sample pattern obtained in the step (g) with a plurality of template patterns, the plurality of template patterns comprising the first and second

template patterns and the template patterns of No. 3 to No. n, n pieces of template patterns obtained in the steps (b), (c) and (e), to find a template pattern substantially identical to the sample pattern; and

- (i) determining the base sequence of the target single-stranded nucleic acid to be a base sequence complementary to the base sequence of the probe taken for the preparation of the identified template pattern found in the step (h).
- 3. (Original) The method according to claim 2, wherein the step (g) further comprises the substep of obtaining a two-valued pattern of the fluorescence intensity by using the threshold fluorescence intensity Fi.
- 4. (Original) The method according to claim 2, wherein the length of the probe is 8 mer to 30 mer.
- 5. (Original) The method according to claim 4, wherein the length of the probe is 12 mer to 25 mer.
- 6. (Original) The method according to claim 2, wherein the number of the mismatched base pairs (i) is 1.
 - 7. and 8. (Cancelled)

- 9. (Currently Amended) A method for identifying a mutation at a plurality of sites in a region in a target single-stranded nucleic acid comprising the steps of:
- (a) preparing a probe array in which single-stranded nucleic acid probes of No. 1 to No. n ($n \ge 3$) are arranged as isolated spots on a substrate, where probes as a whole cover every mutation expected at the sites;
- (b) reacting a single-stranded nucleic acid, which has a base sequence fully complementary to a base sequence of one of the single-stranded nucleic acid probes and is fluorescence-labeled, with the probe array under such conditions that single-stranded nucleic acids complementary to each other form a double-stranded nucleic acid;

removing the unreacted labeled single-stranded nucleic acid; and
measuring fluorescence intensity of each spot of the probe array to obtain a
first image template pattern showing a relationship between location of the probes and
fluorescent characteristics:

- (c) performing the same operation as the step (b) for each of remaining single-stranded nucleic acid probes using a second to a nth single-stranded nucleic acid, and obtaining image template patterns of No. 2 to No. n showing a relationship between location and fluorescent characteristics of the probes;
- (d) performing the same operation as the step (b) using a sample containing the target single-stranded nucleic acid of the base sequence to obtain a sample pattern showing a relationship between a position and fluorescent characteristics;
- (e) comparing the sample pattern obtained in the step (d) with a plurality of image template patterns, the plurality of image template patterns comprising the first image

template pattern and the image template patterns of No. 2 to No. n, to find an image n pieces of said image template patterns obtained in the steps (b) and (c), to find a template pattern substantially identical to the sample pattern; and

(f) identifying the mutation in the target nucleic acid from the sequence of the single-stranded nucleic acid that produced the substantially identical image template pattern found in the step (e).